### **PATENT COOPERATION TREATY**

From the INTERNATIONAL SEARCHING AUTHORITY

To:					PCT	
		•				
	··· see form	PCT/ISA/220			TEN OPINION OF THE	
	300 101111	1 01/10/4220		INTERNATION	NAL SEARCHING AUTHORITY	
				(F	PCT Rule 43bis.1)	
				Date of mailing		
			. <u></u>	(day/month/year) see	e form PCT/ISA/210 (second sheet)	
	licant's or agent's file			FOR FURTHER A	ACTION	
see	form PCT/ISA/2	220		See paragraph 2 below		
i _	mational application		International filing date (d	lay/month/year)	Priority date (day/month/year)	
	T/CA2004/00064	-	30.04.2004		30.04.2003	
			both national dassification a		4045040	
		14/50, A6 IN36/	33, A61K38/18, C12N	15/09, AU1K6//02/,	A61P3/10	
	licant .RATAH PHARM	MACEUTICAL S	INC			
			, INC.	<u> </u>		
1.	This opinion co	ontains indication	ons relating to the folio	owing items:		
	Box No. I	Basis of the op	inion	•		
	Box No. II	Priority				
	⊠ Box No. III			rd to novelty, inventive	e step and industrial applicability	
	☐ Box No. IV	Lack of unity of			•	
	Box No. V	Reasoned state applicability; cit	ement under Rule 43 <i>bis.</i> tations and explanations	1(a)(i) with regard to r supporting such state	novelty, inventive step or industrial ement	
	Box No. VI	Certain docum	ents cited			
	☐ Box No. VII		in the international appl			
	☐ Box No. VIII	Certain observ	ations on the internation	al application	i	
2.	FURTHER ACT	ION			~	
	If a demand for international preliminary examination is made, this opinion will usually be considered to be a written opinion of the International Preliminary Examining Authority ("IPEA"). However, this does not apply where the applicant chooses an Authority other than this one to be the IPEA and the chosen IPEA has notifed the International Bureau under Rule 66.1 bis(b) that written opinions of this International Searching Authority will not be so considered.					
	submit to the IPt	:A a written reply date of mailing o	together, where approp	riate, with amendmen	PEA, the applicant is invited to its, before the expiration of three if 22 months from the priority date,	
	For further option	ns, see Form PC	T/ISA/220.			
3.	For further detail	s, see notes to F	form PCT/ISA/220.			
					!	
					!	

Name and mailing address of the ISA:

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Fayos, C

**Authorized Officer** 

Telephone No. +49 89 2399-2180



International application No. PCT/CA2004/000648

-	D. C.	- No. 1 - Do to the state of th				
-		c No. I Basis of the opinion				
1.	<ol> <li>With regard to the language, this opinion has been established on the basis of the international application in the language in which it was field, unless otherwise indicated under this item.</li> </ol>					
		This opinion has been established on the basis of a translation from the original language into the following language , which is the language of a translation furnished for the purposes of international search (under Rules 12.3 and 23.1(b)).				
2.	<ol> <li>With regard to any nucleotide and it amino acid sequence disclosed in the international application and necessary to the claimed invention, this opinion has been established on the basis of:</li> </ol>					
	a. type of material:					
	Σ	a sequence listing				
	C	able(s) related to the sequence listing				
	b. format of material:					
	×	in written format				
	×	in computer readable form				
	c. time of filing/furnishing:					
	×	contained in the international application as filed.				
		filed together with the international application in computer readable form.				
	Ø					
3.	h	n addition, in the case that more than one version or copy of a sequence listing and/or table relating thereto as been filed or furnished, the required statements that the information in the subsequent or additional opies is identical to that in the application as filed or does not go beyond the application as filed, as ppropriate, were furnished.				
4.	Additional comments:					

International application No. PCT/CA2004/000648

_	RO	x No. II	Priority				
1.	Ø	The fol	lowing document has not been furnished:				
		×	copy of the earlier application whose priority has been claimed (Rule 43bis.1 and 66.7(a)).				
			translation of the earlier application whose priority has been claimed (Rule 43bis.1 and 66.7(b)).				
		Consec	quently it has not been possible to consider the validity of the priority claim. This opinion has neless been established on the assumption that the relevant date is the claimed priority date.				
2.		This op	ninion has been established as if no priority had been claimed due to the fact that the priority claim en found invalid (Rules 43 <i>bis</i> .1 and 64.1). Thus for the purposes of this opinion, the international ate indicated above is considered to be the relevant date.				
3.	Add		bservations, if necessary:				

International application No. PCT/CA2004/000648

Box No. III Non-establishment of opinion with regard to novelty, inventive step and industrial applicability							
Ti ol	The questions whether the claimed invention appears to be novel, to involve an inventive step (to be non obvious), or to be industrially applicable have not been examined in respect of:						
. 0							
Ø	claims Nos. 1-68 completely; claims 23-28, 31-36, 40-62 industrial applicability in particular						
be	because:						
⋈	the said international application, or the said claims Nos. claims 23-28, 31-36, 40-62 industrial applicability relate to the following subject matter which does not require an international preliminary examination (specify):						
	see separate sheet						
Ø	the description, claims or drawings (indicate particular elements below) or said claims Nos. 1-68 are so unclear that no meaningful opinion could be formed (specify):						
	see separate sheet						
⊠	the claims, or said claims Nos. 1-68 are so inadequately supported by the description that no meaningful opinion could be formed.						
	no international search report has been established for the whole application or for said claims Nos.						
	the nucleotide and/or amino acid sequence listing does not comply with the standard provided for in Annex C of the Administrative Instructions in that:						
	the written form		has not been furnished				
			does not comply with the standard				
	the computer readable form		has not been furnished				
			does not comply with the standard				
	the tables related to the nucleo not comply with the technical re	and/or amino acid sequence listing, if in computer readable form only, do ements provided for in Annex C-bis of the Administrative Instructions.					
	See separate sheet for further details						

International application No. PCT/CA2004/000648

Box No. V Reasoned statement under Rule 43*bis*.1(a)(i) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. Statement

Novelty (N)

Yes: Claims

No: Claims

Inventive step (IS)

Yes: Claims

No: Claims

Industrial applicability (IA)

Yes: Claims

23-28, 31-36, 40-62 see separate sheet

No: Claims

2. Citations and explanations

see separate sheet

### Re Item III

Non-establishment of opinion with regard to novelty, inventive step and industrial applicability

- 1- Claims 23-28, 31-36, 40-62 relate to subject-matter considered by this Authority to be covered by the provisions of Rule 67.1(iv) PCT. Consequently, no opinion will be formulated with respect to the industrial applicability of the subject-matter of these claims (Article 34(4)(a)(I) PCT).
- 2- Although claims 1, 23, 24, 29, 30, 31, 32, 33, 34, 35, 36, 37, 40, 41, 42, 44, 56, 59, 60, 63, 67, 68 have been drafted as separate independent claims, they appear to relate effectively to the same subject-matter and to differ from each other only with regard to the definition of the subject-matter for which protection is sought in respect of the terminology used for the features of that subject-matter. The aforementioned claims therefore lack conciseness and as such do not meet the requirements of Article 6 PCT.
- 2.1- Therefore, no opinion is to be formulated on the novelty and inventive step for the subject matter of claims 1-68.
- 3- Furthermore, claims 1-20, 23-68 do not meet the requirements of Article 6 PCT in that the matter for which protection is sought is not clearly defined. The claims attempt to define the subject-matter in terms of the result to be achieved or in an unclear manner (KGF <u>agonist</u>, gastrin <u>compound</u>, gastrin/CCK <u>receptor ligand</u>) which merely amounts to a statement of the underlying problem, without providing the technical features necessary for achieving this result.
- 3.1- In addition, claims 1-68 are not supported by the description as required by Article 6 PCT, as their scope is broader than justified by the description. The description only provides support in terms of technical data for the combination KGF/gastrin (see examples).

If the applicant does not restrict the subject matter of claims 1-68 to the combination KGF/gastrin (Art. 6 PCT), then he should indicate why it is believed that the other claimed, but non-exemplified possibilities are active. It should be borne in mind that a technical effect solving a technical problem has to be achieved by all embodiments falling within the scope of the claim.

- 4.2- Furthermore, since the present application provides one single example of combination (namely gastrin / KGF), claims 1-68 lack clarity, support and disclosure (Arts. 6 and 5 PCT), since the skilled person, after reading the description, would not able to perform the invention over the whole area claimed without undue burden and without needing inventive skill.
  - Indeed, the claims encompass a genus of compounds defined only by their function wherein the relationship between the structural features of the members of the genus and said function have not been defined. In the absence of such a relationship either disclosed in the application as originally filed or which would have been recognised based on information readily available to the skilled man, the skilled person would not know how to make and use compounds that lack any structural definition. It would require undue experimentation (be an undue burden) to randomly screen undefined compounds for the claimed activity.
- 4.3- Hence, no opinion with regards to the novelty, inventive step and industrial applicability is to be formulated for the subject matter of present claims 1-68.

### For the sake of completeness, the following is to be noted:

#### Re Item V

Reasoned statement with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

- 5- The following documents are cited; the relevant passages are those indicated in the search report, unless otherwise specified:
- D1: US-A-5 858 977 (AUKERMAN SHARON LEA ET AL) 12 January 1999 (1999-01-12)
- D2: WO 00/44400 A (RTP PHARMA INC ; GEN HOSPITAL CORP (US)) 3 August 2000 (2000-08-03)
- D3: YAMAOKA T ET AL: "Development of pancreatic islets (review)." INTERNATIONAL JOURNAL OF MOLECULAR MEDICINE. MAR 1999, vol. 3, no. 3, March 1999 (1999-03), pages 247-261, XP009037276 ISSN: 1107-3756
- D4: LOGSDON CRAIG D ET AL: "Adenoviral-mediated gene transfer of dominant negative ras inhibits pancreatic acinar cell growth responses to cholecystokinin and fibroblast growth factor" GASTROENTEROLOGY, vol. 112, no. 4 SUPPL., 1997, page A458, XP009037266 & DIGESTIVE DISEASE WEEK AND THE 97TH ANNUAL

MEETING OF THE AMERICAN GASTROENTEROLOGICAL ASSOCIATION; WASHINGTON, D.C., USA; MAY 11-14, 1997 ISSN: 0016-5085

- 5.1- As mentioned above, no opinion will be formulated with regards to the novelty and inventive step for the subject matter of claims 1-68. It should be noted however, that the combination of KGF and gastrin for the treatment of diabetes and related diseases appears obvious from D1, taken in combination with D2.
- 6- For the assessment of the present claims 23-28, 31-36, 40-62 on the question whether they are industrially applicable, no unified criteria exist in the PCT Contracting States. The patentability can also be dependent upon the formulation of the claims. The EPO, for example, does not recognize as industrially applicable the subject-matter of claims to the use of a compound in medical treatment, but may allow, however, claims to a known compound for first use in medical treatment and the use of such a compound for the manufacture of a medicament for a new medical treatment.
- 7- When / if carrying out amendments, and in order to facilitate the examination of the conformity of the amended application with the requirements of Article 34(2)(b) PCT, the applicant is requested to clearly identify the amendments carried out, no matter wether they concern amendments by addition, replacement or deletion, and to indicate precisely the passages of the application as filed on which these amendments are based (also rule 66.8 (a) PCT).

Only amendments with a clearly identified basis on the application as originally filed will be taken into account for the international preliminary examination report.